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### (54) LYMPHOCYTE ACTIVATION ANTIGEN HB15, A MEMBER OF THE IMMUNOGLOBULIN SUPERFAMILY

LYMPHOZYTENAKTIVIERUNGSANTIGEN HB15 EIN MITGLIED DER IMMUNGLOBULIN SUPERFAMILIE

ANTIGENE HB15 D'ACTIVATION LYMPHOCYTAIRE APPARTENANT AU SOUS-ORDRE DES IMMUNOGLOBULINES

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- EMBL Database entry EBV, accession no. V01555 et al.; 1983; Epstein-Barr virus genome.
- JOURNAL OF IMMUNOLOGY vol. 143, 1989, pages 712 - 717 T.F. TEDDER ET AL.; 'Isolation of cDNAs encoding the CD19 antigen of human and mouse B lymphocytes' cited in the application
- JOURNAL OF IMMUNOLOGY vol. 149, 1992, pages 735 - 742 L.-J. ZHOU ET AL.; 'A novel cell-surface molecule expressed by human interdigitating reticulum cells, Langerhans cells, and activated lymphocytes is a new member of the Ig superfamily'

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**Description**

[0001] This invention relates to nucleic acid sequences encoding a human lymphocyte activation antigen termed HB15, and to the proteins and polypeptides encoded by those sequences.

[0002] Many of the cell-surface molecules which regulate immune responses contain conserved structural features similar to those found in immunoglobulin (Ig). These molecules are encoded by genes that are presumed to have evolved from a common precursor and are therefore members of a large superfamily (Williams et al., *Annu. Rev. Immunol.*, **7**: 381-405 (1988)). Many of the Ig superfamily members are involved in cell-cell adhesion and signal transduction. While most members of this family contain multiple linearly-assembled Ig-like domains, several proteins have been identified that contain single Ig-like domains. Single Ig-like domain proteins that are known or assumed to be involved in cell-cell adhesion include: CD6 $\alpha$  (Litman et al., *Cell* **40**: 237 (1985)), CD8 $\beta$  (Johnson et al., *Nature* **323**: 74 (1986)), CD7 (Aruffo et al., *EMBO J.* **6**: 3313 (1987)), Thy-1 (Williams et al., *Science* **216**: 696 (1982)), CD28 (Aruffo et al., *Proc. Natl. Acad. Sci. USA* **84**: 6573 (1987)), CTLA-4 (Brunet et al., *Nature* **328**: 267 (1987)) and P0 which is a structural protein of the peripheral myelin sheath (Lemke et al., *Cell* **40**: 501 (1985)). In addition, others associate with the antigen receptors of B and T lymphocytes forming multimeric signal-transducing complexes including: CD3  $\gamma$ ,  $\delta$  and  $\epsilon$  chains (Gold et al., *Nature* **321**: 431-434 (1986); van den Elsen et al., *Nature* **312**: 413-418 (1984)), B29 (Hermanson et al., *Proc. Natl. Acad. Sci. USA* **85**: 6890 (1988)), and mB1 (Sakaguchi et al., *EMBO J.* **7**: 3457-3464 (1988)).

[0003] Two single Ig-like domain containing proteins found on lymphocytes are preferentially associated with cellular activation and are known to be involved in mediating cell-cell interactions. CD28 is expressed much more on activated than nonactivated T and B lymphocytes (Turka et al., *J. Immunol.* **144**: 1646 (1990)), and CTLA-4 is expressed mostly, if not exclusively, by activated T and B lymphocytes (Brunet et al., *Nature* **328**: 267 (1987); Harper et al., *J. Immunol.* **147**: 1037-1044 (1991)). The role of CD28 as a T cell receptor for the B7-molecule expressed by activated B cells has been recently identified (Linsley et al., *Proc. Natl. Acad. Sci. USA* **87**: 5031-503 (1990); Freeman et al., *J. Immunol.* **143**: 2714-2722 (1989)), as has a similar role for CTLA-4 (Linsley et al., *J. Exp. Med.* **174**: 561-569 (1991)). As with CD28 and B7, most of the Ig-like domain-containing receptors interact with other members of the Ig superfamily present on other cells.

[0004] cDNA cloned from a human lymphocyte library has been analyzed and shown by the present inventors to encode a novel cell-surface glycoprotein, termed HB15, expressed by activated lymphocytes. The mature 186 amino acid protein encoded by the cDNA was composed of a single extracellular V type immunoglobulin (Ig)-like domain, a transmembrane domain and a 39 amino acid cytoplasmic domain. Northern blot analysis revealed that HB15 derives from three mRNA transcripts of ~1.7, 2.0 and 2.5 kb expressed by lymphoblastoid cell lines. Monoclonal antibodies reactive with HB15 were produced and used to show that HB15 is expressed as a single chain cell-surface glycoprotein of M, 45,000. HB15 expression was specific for lymphoblastoid cell lines and mitogen-activated lymphocytes; HB15 was not expressed at detectable levels by circulating leukocytes. Immunohistological analysis revealed that HB15 has a unique pattern of expression among tissues, being found predominantly in hematopoietic tissues with scattered expression by interfollicular cells and weak expression by mantle zone and germinal center cells. Uniquely, HB15 is also expressed by Langerhan's cells within the skin and circulating dendritic cells. Thus, the HB15 glycoprotein represents a new member of the Ig superfamily.

[0005] cDNA sequences encoding the HB15 protein or portions thereof, including any of its specific domains, ligand binding fragments or immunospecific fragments, can be incorporated into replicable expression vectors and the vectors transfected into an appropriate host (e.g., a bacterial, yeast, or eucaryotic cell culture). Alternatively, genomic DNA fragments encoding the HB15 protein or portions thereof can be utilized *in situ*. The expressed proteins or polypeptides, or antagonists thereto, can be used to modulate mammalian immune function. Also, the expressed products can be employed as immunogens in order to raise antibodies against HB15 or portions thereof including any of its specific domains or fragments thereof.

[0006] Thus, the invention generally features nucleic acid isolates encoding lymphocyte activation antigen, HB15, or portions thereof including any of its specific domains, ligand binding fragments or immunospecific fragments; the encoded HB15 protein or portions thereof including specific domains, ligand binding fragments and immunospecific fragments; methods of producing HB15 or portions thereof; methods of detecting the presence of HB15 or of an HB15 ligand; methods of identifying or developing antagonists to HB15 or HB15 ligand function; methods of diagnosing or treating a patient suffering from an immunological disorder, methods of identifying or isolating cells that express HB15 or fragments thereof, and antibodies reactive with HB15 or fragments thereof.

[0007] Also featured are derivatives of HB15 having variant amino acid sequences or glycosylation not otherwise found in nature, the nucleic acid isolates encoding such derivatives, and polynucleotide probes capable of hybridizing under stringent conditions to the HB15 gene.

[0008] As used herein the term "antagonist to HB15" includes any agent which interacts with HB15 and interferes with its function, e.g., antibody reactive with HB15 or any ligand which binds to HB15. The term "identify" is intended to include other activities that require identification of an entity, such as isolation or purification. The terms "isolated"

or "essentially purified" refer to a nucleic acid or protein sequence that has been separated or isolated from the environment in which it was prepared or in which it naturally occurs. Such nucleic acid or protein sequences may be in the form of chimeric hybrids, useful for combining the function of our nucleic acid or protein sequences with other species. The term "immunospecific fragment" refers to a fragment of the indicated protein that reacts with antibodies specific for a determinant of the indicated protein.

[0009] The HB15 protein, immunospecific or ligand binding fragments or specific domains thereof, or other antagonists to HB15 that interfere with HB15 function, can be used therapeutically to modify or inhibit the development or progression of an immune response or cellular interaction, or to deliver drugs, toxins, or imaging agents to cells that express HB15. HB15 cDNA can be used to produce these proteins or peptide fragments; to identify nucleic acid molecules encoding related proteins or polypeptides (e.g., homologous polypeptides from related animal species and heterologous molecules from the same species); or to build other new, chimeric molecules having similar function either in transformed cells or in cell free systems. In addition, HB15 cDNA can be used to synthesize antisense oligonucleotides for inhibiting the expression of the HB15 protein. Assays for HB15 function, production or expression by cells are made possible by the development of monoclonal antibodies selectively reactive with the HB15 protein.

[0010] Other features and advantages of the invention will be apparent from the following description of preferred embodiments thereof.

[0011] In the drawings:

- Fig. 1 shows the structure of the HB15 cDNA clone and the location of restriction sites;
- Fig. 2 shows the cDNA nucleotide sequence and the deduced amino acid sequence of HB15;
- Fig. 3 shows a hypothetical model for the structure of the extracellular domain of HB15;
- Figs. 4A and 4B show the immunofluorescence results obtained with three lymphoblastoid cell lines that express HB15; and
- Figs. 5A-5F show immunohistochemical analysis of HB15 expression.

[0012] The lymphocyte activation antigen, HB15, is expressed exclusively by lymphoid tissue and skin Langerhan's cells. Referring to Fig. 1, the structural features of the HB15 protein, predicted from nucleotide sequence derived from multiple cDNA clones, clearly establish it as a new member of the Ig superfamily. The predicted structure of HB15 is that of a typical membrane glycoprotein, with a single extracellular Ig-like domain, a transmembrane domain and an approximately 40 amino acid cytoplasmic domain. It is likely that the entire coding region for HB15 was identified as transfection of cell lines with pHB15 cDNA generated cell surface expression of the protein and the  $M_r$  of the immunoprecipitated protein was similar in both cDNA transfected cells (~45,000) and HB15<sup>+</sup> Raji cells (~40,000). It is also likely that HB15 undergoes extensive post-translational processing as HB15 was expressed as a single chain molecule, yet the determined  $M_r$  was twice the predicted size of the core protein. Since HB15 was also expressed on the surface of cDNA transfected cells, including COS cells, CHO cells, a mouse pre-B cell line and a human erythroleukemia line, it is likely that surface expression is not dependent on expression of other components of a molecular complex as occurs with the Ig-like proteins that associate with the T and B cell antigen receptors.

[0013] Comparison of the HB15 amino acid sequences with other previously identified proteins did not reveal any striking homologies, except the similarity of the extracellular Ig-like domain with other members of the Ig superfamily. The HB15 Ig-like domain contained many of the conserved features found in the V-set of domains as shown in Fig. 2 (Williams et al., Ann. Rev. Immunol. **5**:361-405 (1996)). Based on the homology with Ig domains, HB15 is likely to possess a disulfide bond linking Cys 16 and Cys 88. This would place 71 amino acids between the two Cys residues which is of the appropriate size for V-related domains (Williams et al., *supra*). There is the potential for additional disulfide bond formation between residues at positions 8, 81 and 110 since these Cys are present in the extracellular domain as well. In addition, HB15 has a Cys residue located within the predicted membrane spanning domain at position 144. Cys residues are also located at identical positions in CD36 and CD7, suggesting some functional significance, perhaps as sites for fatty acylation (Kaufman et al., J. Biol. Chem. **259**:7230-7238, (1984); Rose et al., Proc. Natl. Acad. Sci., USA **81**:2050-2054 (1984)). The HB15 cytoplasmic tail is similar in size to that of CD7 (Aruffo et al., EMBO J. **6**:3313 (1987)), but shared no amino acid sequence similarity with known proteins. However, the five Ser/Thr residues within this domain could serve as potential sites of phosphorylation. Thus, HB15 appears to be a newly described lymphocyte cell surface antigen that shares no apparent relatedness with previously described structures.

[0014] The HB15 extracellular domain is different from the typical Ig-like domain in that it is encoded by at least two exons. Analysis of partial genomic DNA sequence revealed that half of the Ig-like domain is encoded by a single exon and the putative membrane spanning domain is also encoded by a distinct exon (Fig. 2). That Ig-like domains can be encoded by more than one exon has been observed for some members of the Ig superfamily, including the P0 protein (Lemke et al., Neuron **1**:73-83 (1988)), CD4 (Litman et al., Nature **325**:453-455 (1987)) and N-CAM (Owens et al., Proc. Natl. Acad. Sci., USA **84**:294-298 (1987)). This finding supports structural analyses which suggested that Ig domains may have arisen from an ancestral half-domain that evolved through duplication and subsequent adjoining.

However, each of the above genes and the HB15 gene contain introns at different locations between the sequences coding for the conserved Cys residues of the disulfide bond (Williams et al., *Annu. Rev. Immunol.* **8**:381-405 (1989)). This finding supports the notion that introns may have been subsequently inserted to interrupt the ancestral Ig-like domain at later points during the evolution of each of these domains.

5 [0015] Expression of HB15 appears to be generally restricted to lymphocytes since two monoclonal antibodies reactive with HB15 failed to detect HB15 on most other hematopoietic cells. HB15 expression may be a late event in lymphocyte development as most thymocytes and circulating lymphocytes did not express detectable levels of cell surface HB15. However, after being activated by mitogens, peripheral lymphocytes expressed maximal levels of cell surface HB15 on days 3 through 5, the period of time during which maximal proliferation occurred. HB15 may be expressed at low levels by monocytes, especially after culture or activation, but the level of expression is low and may just result from Fc receptor mediated antibody attachment. Many T and B cell lines also expressed HB15, but expression was generally at low levels. Interestingly, cell-surface HB15 expression by cell lines was highest during periods of maximal proliferation such as on the first day after the cultures were fed. These results imply that HB15 is important for maximal growth of lymphoblastoid cells or the maximal growth of cells is critical for the expression of this antigen.

10 15 This was consistent with the observation that HB15 was expressed by germinal center cells in hematopoietic tissues. Nevertheless, HB15 expression appeared to be lymphoid tissue restricted as revealed by immunohistological analysis of twenty-two different tissues. The only exception was the finding that skin Langerhan's cells express HB15. This unique pattern of restricted expression, along with the structural analysis of the protein, indicates that HB15 is a newly identified lymphocyte activation antigen.

20 [0016] The structural similarity of HB15 with other members of the Ig superfamily suggests that it may be involved in cellular interactions since Ig-like domains are frequently involved in a variety of homotypic and heterotypic interactions in the immune and nervous systems. These interactions include binding functions that trigger a subsequent event below the cell surface or adhesion. A key functional feature is that homophilic or heterophilic binding usually occurs between Ig-related molecules, and this is often between molecules on opposed membrane surfaces. The structural relatedness of HB15 to these other proteins may imply a role for this lymphocyte activation protein in either homotypic or heterotypic interactions of lymphocytes following activation or other HB15<sup>+</sup> cell types.

25 [0017] It is understood that the particular nucleotide and amino acid sequences disclosed in Fig. 2 are representative of the counterpart and related human genes and proteins that can conveniently and directly be obtained following the teaching of this disclosure. For example, cross-hybridization under stringent conditions of the disclosed nucleic acid sequences with genetic material from human cells, can readily be performed to obtain equivalent human sequences. In an analogous manner, degenerate oligonucleotides can readily be synthesized from the disclosed amino acid sequence, or portions thereof, and amplified using any well-known amplification technique, such as the polymerase chain reaction, to obtain probes that bind to equivalent human sequences. Proteins or polypeptides encoded by equivalent sequences can be produced. Antibodies directed against the disclosed protein or peptides can also be raised and employed to cross-react with human and other mammalian peptides having similar epitope(s). Those peptides isolated in this manner that have similar antibody reactivity patterns to those of the disclosed proteins or peptides are considered equivalents of the disclosed proteins or peptides.

30 35 [0018] The following examples are presented to illustrate the advantages of the present invention and to assist one of ordinary skill in making and using the same. These examples are not intended in any way otherwise to limit the scope of the disclosure.

#### EXAMPLE I

##### Isolation and characterization of HB15 cDNA clones and characterization of the HB15 protein

45 [0019] A human tonsil cDNA library was screened by differential hybridization using labeled cDNA from the B lymphoblastoid cell line Raji and the T cell line H-SB2. Two of the 261 RAJI<sup>+</sup> H-SB2<sup>-</sup> cDNA clones isolated, pB10 (~2.5 kb) and pB123 (~1.2 kb), cross hybridized, yet failed to hybridize with cDNA that encode known B cell surface antigens (Tedder et al., *Proc. Natl. Acad. Sci., USA* **85**:208 (1988)). Expression of this mRNA was examined by Northern blot analysis using poly(A)<sup>+</sup> RNA isolated from B cell lines (NALM-6, Narmalwa, Daudi, SB, and Raji), T cell lines (Hut-78, H-SB2, and MOLT-3) and the erythroleukemia line, K562. The pB123 cDNA hybridized strongly with three mRNA species of ~1.7, ~2.0 and ~2.5 kb in SB and Raji, Daudi and Narmalwa cells expressed lower levels of this mRNA. Further autoradiography of the blot (7 days) revealed that the NALM-6, Hut-78 and MOLT-3 cells also expressed these three mRNA species, but at much lower levels, and faint hybridization with H-SB2 RNA was detected. These results suggested differential expression of this gene among leukocyte subpopulations.

50 55 [0020] Restriction maps were generated for these cDNA and their nucleotide sequences determined. Both cDNA were overlapping and contained open reading frames at their 5' ends with the pB123 cDNA having the longest 5' sequence. Since neither clone contained a translation initiation site, the pB10 cDNA insert was used to isolate 13

additional cross-hybridizing cDNA from a human tonsil library. Restriction maps and nucleotide sequence determination indicated that 12 of the cDNA were overlapping, with one cDNA having the longest sequence at the 5' end. The restriction map and nucleotide sequence of this clone, termed pHB15, is shown in Fig. 1. The full length cDNA clone is likely to include an ~500 bp fragment at the 3' end that was removed from the cDNA by EcoRI digestion and subcloning.

5 Eight other independent cDNA clones had similar EcoRI generated fragments and an EcoR I site was located at the identical nucleotide position in all cDNA that were sequenced.

[0021] The pHB15 cDNA had a 625 bp open reading frame, with the major portion of the cDNA representing untranslated sequence. The determined nucleotide sequence and predicted amino acid sequence of HB15 are given in Fig. 2. The predicted cleavage site used to generate the mature protein is shown by a vertical arrow. The numbers shown above the amino acid sequence designate amino acid residue positions of the putative mature protein and the numbers on the right designate nucleotide positions. Amino acids are designated by the single-letter code, and \* indicates the termination codon. Nucleotides delineating translated regions with hydrophobic character are underlined. Amino acids indicating potential N-linked glycosylation attachment sites are underlined. A poly(A) attachment signal sequence is indicated by wavy underlining. The Cys residues are circled and amino acids which are often conserved in Ig-like domains are indicated by (+). Arrow heads below the nucleotide sequence denote exon/intron boundaries identified in another DNA clone.

[0022] The first ATG shown is the most likely initiation codon for translation since it conforms to the proposed translation initiation consensus sequence, (A/G)CCAUG (Kozak, *Cell* **44**:283-292 (1986)). It is likely that the different mRNA species result from differential use of poly(A) attachment sites, AATAAA, since one was found at nucleotide position 248 in the middle of the 3' untranslated region (Fig. 2). This poly(A) attachment site was functional in the pb123 cDNA since it was followed by a poly(A) tail. A poly(A) attachment site or tail was not found in the ~550 bp EcoR I fragment which presumably represents the 3' end of the pHB15 cDNA.

[0023] One clone isolated from the cDNA library (~3.0 kb long) that hybridized with the pb123 cDNA had a unique sequence with 229 and 107 bp long segments that were identical to those found in the other cDNA. These regions had flanking sequences that corresponded to the consensus 5' and 3' splice sequences which demarcate exon boundaries (Aebi et al., *Trends Genet.* **3**:102-107 (1987)) indicating that this aberrant cDNA was composed of introns and two exons. The three splice junction sites identified by this clone are shown (Fig. 2).

[0024] The predicted length of the HB15 protein was 205 amino acids (Fig. 2). However, the pb123 cDNA was missing the codon AAG at nucleotide position 500 so the protein may be one amino acid shorter in some cases. This may result from differential splicing at an exon/intron border, that results in the inclusion or loss of a codon since this codon abuts a potential splice site. A similar phenomenon has been found in the CD19 gene which also encodes a member of the Ig superfamily (Zhou et al., *Immunogenetics* **35**:102-111 (1992)). Hydrophathy analysis of the HB15 amino acid sequence by the method of Kyte et al., *J. Mol. Biol.* **157**:105 (1982) revealed two regions of strong hydrophobicity. The first hydrophobic stretch of 19 amino acids represents a typical signal peptide at the amino terminal end of the protein. The algorithm of von Heijne, *Nucleic Acids Res.* **14**:4683-4690 (1986) predicts that the most probable amino-terminus of the mature protein would be the  $\beta$  following amino acid 19. The second hydrophobic region of 22 amino acids most probably represents the transmembrane region. Three potential N-linked glycosylation attachment sites (N-X-S/T) were found in the extracellular domain. Therefore, the predicted molecular mass of the core protein would be ~20,500.

[0025] Six Cys residues were found in the extracellular domain of HB15 and one in the putative membrane spanning domain. One pair of these residues at positions 16 and 88 delineate Ig-like domains (Williams et al., *Annu. Rev. Immunol.* **8**:381-405 (1989)). This domain contained many of the hallmark amino acids which define the V set of Ig-like domains. A computer search of protein sequences using the Protein Identification Resource Protein Sequence Database showed that no proteins shared significant sequence homology with HB15 other than some members of the Ig superfamily.

[0026] Referring to Fig. 3, a hypothetical model is given for the structure of the extracellular domain of HB15 based on the proposed arrangement of the  $\beta$ -pleated sheets for the V domain of Ig heavy chain. Cys residues are represented as filled circles and amino acids encoded by different exons are indicated by alternatively shaded circles. Numbers represent the predicted amino acid residue positions as in Fig. 2.

## EXAMPLE II

### Production of monoclonal antibodies reactive with HB15.

[0027] Hybridomas were generated by the fusion of NS-1 myeloma cells with spleen cells obtained from mice immunized with pHB15 cDNA-transfected COS cells. Monoclonal antibodies reactive in indirect immunofluorescence assays with HB15 mRNA positive cell lines, but not with HB15 negative cell lines, were isolated. Two of these antibodies, anti-HB15a ( $\text{IgG}_2\text{a}$ ) and anti-HB15b ( $\text{IgG}_3$ ) also reacted with COS cells transfected with the pHB15 cDNA, but did not

react with cells transfected with CD19 cDNA (Tedder et al., J. Immunol. **143**: 712-717 (1989)) or the expression vector alone. In addition, these antibodies reacted with a human erythroleukemia cell line, K562, and a mouse pre-B cell line, 300.19, stably transfected with the pHB15 cDNA. The antibodies did not react with untransfected parent cells, cells transfected with vector alone, or CD19, CD20 (Tedder et al., Proc. Natl. Acad. Sci., USA **85**: 208 (1988)) or LAM-1 (Tedder et al., J. Exp. Med. **170**: 123-133 (1989)) cDNA transfected cells. In all cases, the reactivities of the anti-HB15a and anti-HB15b mAb were identical.

### EXAMPLE III

#### 10 Detection of HB15 expression.

##### Immunoprecipitation of cell surface HB15.

[0028] The anti-HB15a mAb was purified, coupled to beads and used to immunoprecipitate HB15 from detergent solubilized extracts of surface-labeled cell lines. Optimum results were obtained using the K562-HB15 cell line (K562 cells transfected with pHB15 cDNA) since the level of HB15 expression was higher than in other cell lines. The anti-HB15a mAb specifically immunoprecipitated proteins that migrated as a single broad band of ~45,000 M<sub>r</sub>. Similar results were obtained when the immunoprecipitated materials were run under reducing or nonreducing conditions. A similar protein was immunoprecipitated from the Raji cell line except the M<sub>r</sub> was ~40,000. Thus, HB15 was expressed as a noncovalently-associated single chain molecule on the cell surface.

HB15 was expressed by activated lymphocytes.

[0029] The tissue distribution of the HB15 surface antigen was examined by indirect immunofluorescence staining with flow cytometry analysis. Two cell lines that did not express HB15 message were transfected with the pHB15 cDNA subcloned into the Bam HI site of the retroviral vector pZipNeoSV(X). Referring to Fig. 4, the immunofluorescence results obtained with three lymphoblastoid cell lines that express HB15 are demonstrated. The open histograms show the cellular reactivity with the HB15a antibody, and the shaded histograms demonstrate background levels of immunofluorescence staining obtained with unreactive control antibodies. Among 33 cell lines examined, HB15 was expressed at detectable levels by B cell lines (including Raji, Daudi, Namalwa, Arent, BJAB, SB, Jijoy, Akata, and SLA) and T cell lines (including Jurkat, H-9, Rex, H-SB2, and Hut-78). However, HB15 expression was generally low and variable. The highest levels of cell-surface expression were always obtained where the cell cultures were recently split and were thus proliferating maximally. Cell lines that did not express detectable levels of HB-15 included: K562, the B cell lines NALM-6 and Ramos; the T cell lines, MOLT-3, RPMI 8405, PEER, MOLT-14, CEM and HPB-ALL; the myelomonocytic line, HL60; the natural killer cell line, YT; the colon carcinoma lines, Colo-205 and HT29; the lung cell lines, NCI-H69, and NCI-H82; the prostate line, PC3; the melanoma line, MEWO; and the breast tumor lines, ZR75.1, MCF7 and BT20.

[0030] Expression of HB15 by normal blood leukocytes was also examined. However, cell-surface expression of HB15 was not detected at significant levels on circulating lymphocytes, natural killer cells or monocytes in 15 blood samples. Therefore, the possibility that HB15 was expressed following cellular activation was examined by inducing T lymphocyte proliferation with the mitogens concanavalin A (ConA), pokeweed mitogen, phytohemagglutinin-P or phorbol esters (PMA). Expression of HB15 was examined 2, 8, 12, 24, 48, 72, 120 and 240 hours following the initiation of cultures. Appearance of HB15 expression paralleled cellular proliferation such that optimal expression was on days 3 through 5 following the initiation of cultures. Also, the quantity of HB15 expression induced was not correlated with any specific mitogen, but correlated more with the strength of the mitogenic signal such that cell-surface expression was predominantly found on the larger blast cells. Therefore, HB15 was expressed by lymphocytes following activation.

##### Immunohistological analysis of HB15 expression.

[0031] The lymphocyte specificity and tissue distribution of HB15 was also examined by immunohistological analysis of different human tissues. Basically, the anti-HB15a mAb was used to stain thymus, tonsil, spleen, lymph node, kidney, renal pelvis and ureter, Fallopian tube, liver, pancreas, stomach, breast, lung, esophagus, skeletal muscle, skin, uterus, salivary gland, thyroid gland, adrenal gland, heart, appendix and colon. (Referring to Figs. 5A-5F), in most cases, HB15 expression appeared lymphocyte specific in that no significant reactivity was observed in non-lymphoid tissues. Among tonsil and lymph nodes (Fig. 5A), HB15 was expressed reasonably strongly by scattered cells in intrafollicular regions (T cell zones) (Fig. 5C). Although some of these cells may have been lymphoblasts, most were interdigitating reticulum cells (a subpopulation of dendritic cells) since they appeared larger than resting lymphocytes and expressed the CD1 surface molecule (Fig. 5D). Also, some cells (50-80%) within germinal centers (GC; Figs. 5A and 5B) and follicular mantle zones (FM; Fig. 5A), with the morphology of lymphocytes, were weakly HB15<sup>+</sup>. Among spleen, the HB15<sup>+</sup> cells

were predominantly restricted to the white pulp, whereas the red pulp remained largely negative. Again, those large, scattered positive cells in the white pulp are likely to be interdigitating reticulum cells or lymphoblasts. Cortical thymocytes were HB15 negative, while a small subpopulation of medullary cells, presumably thymocytes, was positive (Fig. 5E). Unlike other non-hematopoietic tissues, analysis of skin revealed that some cells with the characteristic scattered branching morphology of Langerhan's cells (a subpopulation of dendritic cells) expressed HB15 at detectable levels (Fig. 5F). Among all non-hematopoietic tissues, where inflammatory infiltrations were apparent, a few scattered lymphocytes were found to express HB15. It is also likely that circulating dendritic cells are HB15<sup>+</sup>, but because of their low frequency they were not readily detected. Similarly, it is also likely that the malignant counterparts of dendritic cells express HB15 and that this molecule can be used as a diagnostic marker for malignant cells as the L428 cell line, which is a neoplastic cell line that was derived from Hodgkin's disease and may represent interdigitating reticulum cells (Schaadt et al., Int. J. Cancer 26:723-731 (1980)), is HB15 positive.

#### Experimental Procedures

##### Isolation of cDNA clones.

[0032] The isolation of cDNA clones by differential hybridization has been described (Tedder et al., Mol. Immunol. 25:1321-1330 (1988)). One clone, pB123, was purified, labeled by nick translation (Fligby et al., J. Mol. Biol. 113: 237-251 (1977)) and used to isolate homologous cDNA by again screening the same human tonsil cDNA library in λgt11 (Wells et al., Proc. Natl. Acad. Sci., USA 83:5639-5643 (1986)) as described (Zhou et al., Immunogenetics 35: 102-111 (1992)). Positive plaques were isolated, cloned and the cDNA inserts were removed by EcoR I digestion and subcloned into pSP65 (Melton et al., Nucleic Acids Res. 12:7035-7056 (1984)). Restriction maps were generated as described by Maniatis et al., Molecular Cloning: A Laboratory Manual, (1982) and nucleotide sequences were determined using the method of Sanger et al., Proc. Natl. Acad. Sci., USA 74:5463-5467 (1977).

[0033] A computer search of nucleotide and protein sequences was conducted using the Protein Identification Resource Data (GenBank release 66 and Swiss-Prot-16). Gap penalties of -1 were assessed during sequence homology analysis for each nucleotide or amino acid in the sequence where a gap or deletion occurred.

##### RNA blot analysis.

[0034] Poly(A)<sup>+</sup> RNA was isolated as described (Maniatis et al., Molecular Cloning: A Laboratory Manual, (1982)). For Northern-blot analysis, 2 μg of poly(A)<sup>+</sup> RNA was denatured with glyoxal, fractionated by electrophoresis through a 1.1% agarose gel and transferred to nitrocellulose (Thomas, Method Enzymol. 100:255 (1983)). The pB123 cDNA insert used as probe was isolated, nick-translated (Fligby et al., J. Mol. Biol. 113:237-251 (1977)) and hybridized with the filters as described (Wahl et al., Proc. Natl. Acad. Sci., USA 76:3683-3687 (1979)). Hybridization at high stringency was with 50% (w/v) formamide, 4X SSC, 10% (w/v) Na dextran sulfate at 42°C. The filters were washed at 65°C with 0.2X SSC, 0.1% SDS. RNA size was determined by comparison with 28S and 18S ribosomal RNA run on the same gels as standards. The same blot was also hybridized with cDNA clones containing a housekeeping mRNA of unknown identity revealing that all mRNA were intact and were similar in quantity of this expressed mRNA. For hybridization at low stringency the conditions are overnight incubation at 42°C in a solution comprising: 20% formamide, 5XSSC (150 mM NaCl, 15 mM trisodium citrate), 50 mM sodium phosphate (pH 7.6), 5X Denhardt's solution, 10% dextran sulfate, and 20 μg/ml denatured, sheared salmon sperm DNA.

##### Cells.

[0035] Human blood was obtained by protocols approved by the Human Protection Committee of Dana-Farber Cancer Institute and mononuclear cells were isolated by Ficoll-Hypaque density gradient centrifugation. Mononuclear cells (10<sup>6</sup>/ml) in complete media (RPMI-1640 supplemented with 15% fetal calf serum, antibiotics and glutamine) were stimulated with phytohemagglutinin-P (2 μg/ml; Difco, Detroit, MI), Con A (10 μg/ml, Miles Laboratories, Elkhart, IN), pokeweed mitogen (10 μg/ml, Gibco/BRL, Bethesda, MD) or phorbol myristate 13-acetate (PMA, 10 ng/ml, Sigma, St Louis, MO) as described (Tedder et al., J. Immunol. 144:532-540 (1990)). Lymphocytes were harvested at the indicated time points, washed once in complete media, and aliquoted for immediate immunofluorescence staining as described below.

[0036] COS cells were transfected with the pHB15 cDNA insert subcloned into a modified CDM8 vector (Aruffo et al., EMBO J. 6:3313 (1987); Tedder et al., J. Immunol. 143:712-717 (1989)) using the DEAE-dextran method as described (Aruffo et al., EMBO J. 6:3313 (1987)). Cell surface expression was examined after 48 hours by indirect immunofluorescence. Stable cDNA transfected cells were produced using the pHB15 cDNA cloned into the BamH I site of the retroviral vector pZipNeoSV(X) in the correct orientation (Cepko et al., Cell 37:1053-1062 (1984)). The murine

pre-B cell line, 300.19, and the human erythroleukemia cell line, K562, were transfected with this vector by electroporation with subsequent selection of stable transfectants using G418 (Gibco/BRL). Cells expressing HB15 were further enriched by reacting the cells with monoclonal antibodies with the subsequent isolation of HB15<sup>+</sup> cells by panning on anti-mouse Ig coated plates.

[0037] Cell lines were grown in RPMI 1640 medium containing 10% fetal calf serum and antibiotics. Cultures of all cell lines were split the day before analysis and were in logarithmic growth.

#### mAb production

[0038] Anti-HB15 mAb were generated as described (Tedder et al., J. Immunol. **144**:532-540 (1990)) by the fusion of NS-1 myeloma cells with spleen cells from BALB/c mice that were repeatedly immunized with COS cells transfected with the HB15 cDNA. Each hybridoma was cloned twice and used to generate ascites fluid. The isotypes of the mAb were determined using a mouse monoclonal antibody isotyping kit from Amronsham (Arlington Heights, IL).

#### Immunofluorescence analysis.

[0039] Cells were kept at 4°C and were examined immediately after isolation. Indirect immunofluorescence analysis of viable cells was carried out after washing the cells three times. The cells were then incubated for 20 min on ice with each mAb as ascites fluid diluted to the optimal concentration for immunostaining. Isotype-matched murine antibodies that were unreactive with human leukocytes were used as negative controls. After washing, the cells were treated for 20 min at 4°C with fluorescein isothiocyanate-conjugated goat anti-mouse Ig antibodies (Southern Biotechnology Associates, Birmingham, AL). Single color immunofluorescence analysis was performed on an Epics Profile flow cytometer (Coulter Electronics, Hialeah, FL). Ten thousand cells were analyzed for each sample.

#### Immunoprecipitation analysis.

[0040] Cells were washed twice, resuspended in saline and labeled by the iodogen method as described (Thompson et al., Biochem. **26**:743-750 (1987)). After washing, the cells were lysed in 1 ml of buffer containing 1% (v/v) Triton X-100 and protease inhibitors as described (Tedder et al., Proc. Natl. Acad. Sci., USA **85**:208 (1988)). Immunoprecipitations were carried out using anti-HB15a mAb or mouse Ig (as a negative control) directly conjugated to Affigel (BioRad, Richmond, VA) at 2 mg of mAb per ml of gel according to the manufacturer's instructions. Cell lysates were precleared twice for 2 hours using 50 µl (50% v/v) of murine Ig coated beads at 4°C. Cell lysates were precleared again overnight.

[0041] Half of the precleared lysate was then incubated with 25 µl of anti-HB15a mAb-coated beads or murine Ig-coated beads with constant rotation at 4°C for 18 hours. Immunoprecipitates were washed and analyzed by SDS-PAGE as described (Tedder et al., Proc. Natl. Acad. Sci., USA **85**:208 (1988)) with half of the sample run in the presence of 5% 2-mercaptoethanol (reducing conditions). M<sub>r</sub> were determined using pre-stained standard molecular weight markers (Gibco/BRL).

#### Immunohistochemistry.

[0042] All tissues were stained applying a modification of the APAAP procedure as described by Cordell et al., J. Histochem. Cytochem. **31**:219-229 (1984). Basically, the slides were first incubated with monoclonal antibody followed by an incubation step with rabbit anti-mouse (bridging) antibody. Subsequently, a monoclonal antibody against alkaline phosphatase pre-incubated with alkaline phosphatase was applied. In order to enhance the sensitivity of this procedure, the number of phosphatase molecules on the surface was increased by using one or two layers of bridging antibody and anti-phosphatase antibody. Bound phosphatase molecules were visualized using new fuchsin as a substrate (Cordell et al., J. Histochem. Cytochem. **31**:219-229 (1984)).

[0043] The HB15 protein or immunospecific fragments thereof, or antibodies or other antagonists to HB15 function, can be used to diagnose or treat a variety of immunological disorders, diseases or syndromes. For such purposes, the soluble external domain would often be employed, typically but not necessarily, polymerized in a multivalent state using, e.g., dextran or polyamino acid carriers or fusion proteins of HB15 fragments and carrier molecules. Liposomes may alternatively be employed as the therapeutic vehicle, in which case the transmembrane domain and preferably at least some of the cytoplasmic domain will also be included.

[0044] For example, since Langerhans' cells are the primary immunocompetent cell in the skin, playing a role in the presentation of antigen to T cells and the induction of contact hypersensitivity, and since HB15 is expressed by Langerhans' cells and may be involved in antigen presentation, it is likely to be involved in the pathogenesis of human skin disease such as psoriasis, autoimmune disorders, organ transplant and AIDS.

[0045] Therefore, antagonists to HB15 function can provide important therapeutic agents for treatment of these dis-

ees. Similarly, since HB15 may serve as an accessory molecule for lymphocyte activation, the HB15 antigen, fragments or domains thereof, may be used as agonists that would augment an immune response.

[0046] More specifically, the dendritic cell is a primary target of the human immunodeficiency virus, the causative agent of AIDS. It has recently been proposed that 80% of AIDS virus *in vivo* is produced by dendrite cells, particularly by Langerhans' cells, circulating dendrite cells and interdigitating reticulum cells (Langhoff et al., Proc. Natl. Acad. Sci. USA 88:7998-8002 (1991)). Also, most infections occur through mucosal surfaces where it is thought that dendrite cells are first infected. Therefore, this reagent provides us with a critical tool for the potential prevention or treatment of AIDS or AIDS related disorders.

[0047] For monitoring certain clinical conditions, it may be advisable to quantitate the levels of endogenous soluble HB15 in a patient's blood serum. Based on the finding that several receptors are now known to be shed during various normal and pathological conditions, it is possible that HB15 is also lost from the cell surface by an enzymatic process. Also, quantitative detection can be useful in a method of identifying leukocytes with abnormal or decreased expression of HB15 for diagnosis and/or detection of leukocyte activation or altered leukocyte function. Additionally, the ability to quantitate the amount of receptor, or fragment thereof, produced during the manufacture of a recombinant therapeutic agent will be advantageous. Quantitation of HB15 levels can be carried out using a number of assay methods known to those of ordinary skill in the art, including an enzyme-linked immunocassay using the monoclonal antibodies that have been produced against HB15.

[0048] Similarly, in treating certain clinical conditions, it may be advisable to remove endogenous soluble HB15 or HB15<sup>-</sup> cells from a patient's blood. This can be done with existing on-line and off-line techniques by employing immunoselection columns containing antibodies or other binding agents directed against the disclosed external domain of HB15.

[0049] There are at present no specific markers for non-follicular dendrite cells in humans. Use of HB15 monoclonal antibody to identify HB15<sup>-</sup> cells now permits the isolation and purification of cells expressing this protein from a population of unrelated cells.

[0050] The HB15 mAb will also be useful for the evaluation and diagnosis of interdigitating cell sarcomas or other malignant cell types expressing this antigen. Therefore, HB15-based agents may be suitable for immunotherapy or immunomaging.

[0051] In addition, assays for HB15 function can be used in further research on the physiological role of this receptor. For example, in preliminary experiments T cell proliferation in the mixed lymphocyte reaction, an assay for T cell activation, can be partially inhibited by the presence of anti-HB15 monoclonal antibodies. This functional assay suggests a role for the HB15 molecule on dendrite cells or monocytes in the initiation of T cell function.

[0052] While the present invention has been described in conjunction with a preferred embodiment, one of ordinary skill, after reading the foregoing specification, will be able to effect various changes, substitutions of equivalents, and other alterations to the compositions and methods set forth herein.

[0053] The following hybridomas were deposited on March 17, 1992, with the American Type Culture Collection (ATCC).

Identification	ATCC Designation
Anti-HB15a Hybridoma cell line, HB15a	HB 10987
Anti-HB15b Hybridoma cell line, HB15b	HB 10988

#### SEQUENCE LISTING

[0054]

(1) GENERAL INFORMATION:

(i) APPLICANT: Dana-Farber Cancer Institute, Inc.

(ii) TITLE OF INVENTION: LYMPHOCYTE ACTIVATION ANTIGEN HB15, A MEMBER OF THE IMMUNOGLOBULIN SUPERFAMILY

(iii) NUMBER OF SEQUENCES: 2

(iv) CORRESPONDENCE ADDRESS:

(A) ADDRESSEE: Weingarten, Schurigin, Gagnebin & Hayes

(B) STREET: Ten Post Office Square

(C) CITY: Boston

5 (D) STATE: MA

(E) COUNTRY: USA

10 (F) ZIP: 02109

(v) COMPUTER READABLE FORM:

(A) MEDIUM TYPE: Floppy disk

15 (B) COMPUTER: IBM PC compatible

(C) OPERATING SYSTEM: PC-DOS/MS-DOS

20 (D) SOFTWARE: PatentIn Release #1 0, Version #1.25

(vi) CURRENT APPLICATION DATA:

(A) APPLICATION NUMBER:

25 (B) FILING DATE:

(C) CLASSIFICATION:

30 (vii) PRIOR APPLICATION DATA:

(A) APPLICATION NUMBER: US 07/870,029

(B) FILING DATE: 17-APR-1992

35 (viii) ATTORNEY/AGENT INFORMATION:

(A) NAME: Heine, Holliday C.

40 (B) REGISTRATION NUMBER: 34,346

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45 (A) TELEPHONE: (617) 542-2290

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50 (C) TELEX: 940675

(2) INFORMATION FOR SEQ ID NO 1:

(i) SEQUENCE CHARACTERISTICS:

55 (A) LENGTH: 1762 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: double

(D) TOPOLOGY: linear

5 (ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

10 (iv) ANTI-SENSE: NO

(ix) FEATURE:

(A) NAME/KEY: CDS

15 (B) LOCATION: 11..625

(ix) FEATURE:

(A) NAME/KEY: mat\_peptide

20 (B) LOCATION: 68..622

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

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	GAATTCCGCC ATG TCG CGC GGC CTC CAG CTT CTG CTC CTG AGC TGC GCC Met Ser Arg Gly Leu Gln Leu Leu Leu Ser Cys Ala -19 -15 -10	49
5	TAC AGC CTG GCT CCC GCG ACG CCG GAG GTG AAG GTG GCT TGC TCC GAA Tyr Ser Leu Ala Pro Ala Thr Pro Glu Val Lys Val Ala Cys Ser Glu -5 1 5 10	97
10	GAT GTG GAC TTG CCC TGC ACC GCC CCC TGG GAT CCG CAG GTT CCC TAC Asp Val Asp Leu Pro Cys Thr Ala Pro Trp Asp Pro Gln Val Pro Tyr 15 20 25	145
	ACG GTC TCC TGG GTC AAG TTA TTG GAG GGT GGT GAA GAG AGG ATG GAG Thr Val Ser Trp Val Lys Leu Leu Glu Gly Gly Glu Glu Arg Met Glu 30 35 40	193
15	ACA CCC CAG GAA GAC CAC CTC AGG GGA CAG CAC TAT CAT CAG AAG GGG Thr Pro Gln Glu Asp His Leu Arg Gly Gln His Tyr His Gln Lys Gly 45 50 55	241
20	CAA AAT GGT TCT TTC GAC GCC CCC AAT GAA AGG CCC TAT TCC CTG AAG Gln Asn Gly Ser Phe Asp Ala Pro Asn Glu Arg Pro Tyr Ser Leu Lys 60 65 70	289
	ATC CGA AAC ACT ACC AGC TGC AAC TCG GGG ACA TAC AGG TGC ACT CTG Ile Arg Asn Thr Thr Ser Cys Asn Ser Gly Thr Tyr Arg Cys Thr Leu 75 80 85 90	337
25	CAG GAC CGG GAT GGG CAG AGA AAC CTA ACT GGC AAG GTG ATC TTG AGA Gln Asp Pro Asp Gly Gln Arg Asn Leu Ser Gly Lys Val Ile Leu Arg 95 100 105	385
30	GTG ACA GGA TGC CCT GCA CAG CGT AAA GAA GAG ACT TTT AAG AAA TAC Val Thr Gly Cys Pro Ala Gln Arg Lys Glu Glu Thr Phe Lys Lys Tyr 110 115 120	433
	AGA GCG GAG ATT GTC CTG CTG GCT GTT ATT TTC TAC TTA ACA Arg Ala Glu Ile Val Leu Leu Ala Leu Val Ile Phe Tyr Leu Thr 125 130 135	481
35	CTC ATC ATT TTC ACT TGT AAG TTT GCA CGG CTA CAG ACT ATC TTC CCA Leu Ile Ile Phe Thr Cys Lys Phe Ala Arg Leu Gln Ser Ile Phe Pro 140 145 150	529
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	GAT TTT TCT AAA GCT GGC ATG GAA CGA GCT TTT CTC CCA GTT ACC ACC TCC Asp Phe Ser Lys Ala Gly Met Glu Arg Ala Phe Leu Pro Val Thr Ser 155 160 165 170	577
5	CCA AAT AAC CAT TTA GGG CTA GTG ACT CCT CAC AAG ACA GAA CTG GTA Pro Asn Lys His Leu Gly Leu Val Thr Pro His Lys Thr Glu Leu Val 175 180 185	625
10	TGAGCAGGAT TTCTGCAGGT TCTTCTTCCT GAAGCTGAGG CTCAGGGTG TGCCCTGTC TTACACTGGA GGAGAGAAGA ATGAGCCCTAC GCTGAAGATG GCATCCTGTC AAGTCCTTCA CCTCACTGAA AACATCTGGA AGGGGATCCC ACCCCATTCT CTGTGGGAG GCCTCGAAAA CCATCACATG ACCACATAGC ATGAGGCCAC TGCTGCTTCT CCATGGCCAC CTTTCAGCG 15 ATGTATGCAG CTATCTGGT AACCTCTGG ACATTTTTTC AGTCATATAA AAGCTATGGT GAGATGCAGC TGGAAAAGG TCTTGGAAA TATGAATGCC CCCAGCTGGC CCGTGACAGA 20 CTCTGAGGA CAGCTGCTCT CTCTGCATC TTGGGACAT CTCTTGAAAT TTCTCTGTT TTGCTGTACCC AGCCCATGATGTTTACGTCT GGGAGAAATT GACAGATCAA GCTGTGAGAC AGTGGGAAATT ATTTAGCAA TAATTCCTG CTGTGAAGGT CCTGCTTATAA CTAAAGGAGTA 25 ATCTGTGTAC AAAGAAATATAA CAAGTCGATG AACTATTCCC CAGCAGGGTC TTTTCATCTG GGAAAAGACAT CCATAAAGA GCAATAAAGA AGAGTGCAC ATTATTTTTT ATATCTATAT GTACTTGTCA AAGAAGGTTT GTGTTTTCT GCTTTGAAA TC TGTATCTG TAGTGAGATA 30 GCATTGAACT CTGACAGGCA GCCTGGACAT AGAGAGGGAG AGAAAGTCAG AGAGGGTGC AAGATAGAGA GCTATTAAAT GGCGGGCTG AAATGCTGGG CTGACGGTGC AGTCTGGGTG CTCGTCCACT TGCCCCACTA TCTGGGTGCA TGATCTTGAG CAAGTTCCCTT CTGGTGTCTG 35 CTTTCTCCAT TGTAACACCAC AAGGCTGTG CATGGGCTAA TGAAGATCAT ATACGTGAAA ATTCTTGAA AACATATATAA GCACTATACA GATTGAAAC TCCATTGAGT CATTATCCTT GCTATGATGA TGGTGTGTTTG GGGATGAGAG GGTGCTATCC ATTTCTCATG TTTCCATTG 40 TTGAAACAA AGAAGGTTAC CAAGAACGCT TTCCTGTAGC CTTCTGTAGG AATTCCA 1762	685 745 805 865 925 985 1045 1105 1165 1225 1285 1345 1405 1465 1525 1585 1645 1705

(2) INFORMATION FOR SEQ ID NO:2:

## (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 205 amino acids

(B) TYPE: amino acid

(D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: protein

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Met Ser Arg Gly Leu Gln Leu Leu Leu Ser Cys Ala Tyr Ser Leu  
 -19 -15 -10 -5  
 5 Ala Pro Ala Thr Pro Glu Val Lys Val Ala Cys Ser Glu Asp Val Asp  
 1 5 10  
 Leu Pro Cys Thr Ala Pro Trp Asp Pro Gln Val Pro Tyr Thr Val Ser  
 15 20 25  
 10 Trp Val Lys Leu Leu Glu Gly Gly Glu Arg Met Glu Thr Pro Gln  
 15  
 30 35 40 45  
 15 Glu Asp His Leu Arg Gly Gln His Tyr His Gln Lys Gly Gln Asn Gly  
 50 55 60  
 Ser Phe Asp Ala Pro Asn Glu Arg Pro Tyr Ser Leu Lys Ile Arg Asn  
 65 70 75  
 20 Thr Thr Ser Cys Asn Ser Gly Thr Tyr Arg Cys Thr Leu Gln Asp Pro  
 80 85 90  
 Asp Gly Gln Arg Asn Leu Ser Gly Lys Val Ile Leu Arg Val Thr Gly  
 95 100 105  
 25 Cys Pro Ala Gln Arg Lys Glu Glu Thr Phe Lys Lys Tyr Arg Ala Glu  
 110 115 120 125  
 Ile Val Leu Leu Ala Leu Val Ile Phe Tyr Leu Thr Leu Ile Ile  
 130 135 140  
 30 Phe Thr Cys Lys Phe Ala Arg Leu Gln Ser Ile Phe Pro Asp Phe Ser  
 145 150 155  
 Lys Ala Gly Met Glu Arg Ala Phe Leu Pro Val Thr Ser Pro Asn Lys  
 160 165 170  
 35 His Leu Gly Leu Val Thr Pro His Lys Thr Glu Leu Val  
 175 180 185

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## Claims

1. An isolated nucleic acid comprising a sequence encoding the extracellular domain of the HB15 protein described by SEQ ID NO: 2.
2. The nucleic acid of Claim 1, comprising a sequence encoding the entire HB15 sequence shown in SEQ ID NO:2.
3. The nucleic acid of Claim 1, encoding the amino acid sequence corresponding to residues 1-113 of SEQ ID NO:2.
4. The nucleic acid of Claim 1, encoding the amino acid sequence corresponding to residues 1-125 of SEQ ID NO:2.
5. The nucleic acid of Claim 1, encoding the amino acid sequence shown in SEQ ID NO:2.
6. An isolated nucleic acid comprising a sequence encoding the transmembrane domain of the HB15 protein described by SEQ ID NO:2.
7. The nucleic acid of Claim 6, encoding the amino acid sequence corresponding to residues 126-147 of SEQ ID NO:2.

8. An isolated nucleic acid comprising a sequence encoding the cytoplasmic domain of the HB15 protein described by SEQ ID NO:2.

9. The nucleic acid of Claim 8, encoding the amino acid sequence corresponding to residues 148-186 of SEQ ID NO:2.

10. An isolated nucleic acid comprising a sequence encoding a mammalian homolog of the HB15 protein, wherein said nucleic acid is capable of hybridizing under stringent conditions with a DNA probe comprising the coding sequence shown in SEQ ID NO: 1, and wherein said homolog has the tissue distribution observed for the human HB15 protein described by SEQ ID NO: 2.

11. An isolated nucleic acid comprising that portion of the nucleic acid of Claim 10 which encodes the extracellular domain of said homolog.

12. An isolated nucleic acid comprising that portion of the nucleic acid of Claim 10 which encodes the transmembrane domain of said homolog.

13. An isolated nucleic acid comprising that portion of the nucleic acid of Claim 10 which encodes the cytoplasmic domain of said homolog.

14. A recombinant vector comprising the nucleic acid of any one of Claims 1-13.

15. A cultured cell transfected with the vector of Claim 14.

16. The cell of Claim 15, where said cell in the untransfected form does not express the protein encoded by said nucleic acid.

17. A method of expressing recombinant nucleic acid which comprises culturing the cell of Claim 16 under conditions which allow for the expression of the nucleic acid with which it has been transfected.

18. A method of producing the HB15 protein described by SEQ ID NO:2 comprising incubating the cultured cell of Claim 15 under conditions which allow for production of HB15 protein, and recovering the HB15 protein from the incubated cell.

19. A polynucleotide greater than about 20 nucleotides in length capable of hybridizing under stringent conditions to a nucleic acid having a sequence complementary to a nucleic acid sequence shown in SEQ ID NO:1 encoding the HB15 protein described by SEQ ID NO:2 or a portion thereof having the tissue distribution pattern of HB15.

20. The polynucleotide of Claim 19 wherein the stringent conditions include a hybridization solution comprising 20% formamide, 150mM NaCl, 15mM trisodium citrate, 50mM sodium phosphate, 10% dextran sulphate and 20 µg/ml denatured, sheared salmon sperm DNA.

21. A polynucleotide of Claim 19 that is greater than about 50 nucleotides.

22. A polynucleotide of Claim 21 that is greater than about 100 nucleotides.

23. The polynucleotide of Claim 19, wherein said portion includes an extracellular domain of the HB15 protein described by SEQ ID NO: 2.

24. The polynucleotide of Claim 23 wherein said portion includes amino acids 1-113 of SEQ ID NO:2.

25. A cultured cell transfected with the polynucleotide of Claim 19, operably linked to a transcriptional control sequence.

26. The cell of Claim 25, where said cell in the untransfected form does not express the protein encoded by said polynucleotide.

27. A method of producing the polypeptide encoded by the polynucleotide of claim 19 comprising culturing the cells of Claim 25 under conditions effective for the production of said polypeptide, and recovering said polypeptide.

28. A polypeptide encoded by the nucleic acid of any one of Claims 1-13 and 19-24.

29. A polypeptide having an amino acid sequence of the polypeptide of Claim 28.

5 30. A method of isolating nucleic acid encoding a homologue of the human HB15 protein described by SEQ ID NO: 2, comprising hybridizing said nucleic acid of Claim 1 with a population of nucleic acid molecules suspected of containing a nucleic acid molecule encoding an HB15 homolog under hybridization conditions sufficient to identify said HB15 homolog-encoding nucleic acid molecule within said population by whether hybridization occurs, and isolating said nucleic acid molecule encoding said HB15 homolog from cross-hybridized molecules.

10 31. The method of Claim 30 wherein said population of nucleic acid molecules is obtained from murine tissue.

32. A nucleic acid isolate able to hybridize under stringent hybridization conditions to a nucleic acid as defined by Claim 1.

15 33. A polypeptide encoded by the nucleic acid of Claim 32 and having the tissue distribution pattern of the HB15 protein described by SEQ ID NO: 2.

34. A polypeptide having the sequence of the polypeptide of Claim 33.

20 35. A monoclonal antibody that binds to the polypeptide of Claim 29 or 34.

36. A monoclonal antibody that binds to HB15 protein described by SEQ ID NO: 2.

25 37. A monoclonal antibody that binds to the HB15 epitope recognized by an antibody produced by the cell line deposited as ATCC No: HB10987.

38. A monoclonal antibody that binds to the HB15 epitope recognized by an antibody produced by the cell line deposited as ATCC No: HB10988.

30 39. The monoclonal antibody produced by the cell line deposited as ATCC No: HB10987.

40. The monoclonal antibody produced by the cell line deposited as ATCC No: HB10988.

35 41. The hybridoma cell line deposited as ATCC No: HB10987.

42. The hybridoma cell line deposited as ATCC No: HB10988.

40 43. A method of isolating cells expressing HB15 comprising reacting the antibody of any of Claims 35 to 40 with a population of cells and isolating cells to which said antibody binds.

44. A method of quantitating the amount of surface HB15 expressed on a population of cells, comprising reacting an antibody that binds to the HB15 protein described by SEQ ID NO: 2 with a population of cells, at least some of which are suspected of bearing surface HB15, under conditions which allow said antibody to bind to surface HB15; detecting cells to which said antibody binds; and quantitating the amount of bound antibody.

45 45. The monoclonal antibody of Claim 35, in which the monoclonal antibody is a human monoclonal antibody.

46. The monoclonal antibody of Claim 35, in which the monoclonal antibody is a chimerized monoclonal antibody having variable region segments derived from a mouse monoclonal antibody and other regions derived from a human antibody.

**Patentansprüche**

55 1. Isolierte Nucleinsäure, welche eine die durch SEQ-ID-NR.:2 beschriebene extrazelluläre Domäne des HB15-Proteins kodierende Sequenz enthält.

2. Nucleinsäure nach Anspruch 1, welche eine die in SEQ-ID-NR. 2 dargestellte gesamte HB15-Sequenz kodierende Sequenz enthält.
3. Nucleinsäure nach Anspruch 1, welche die den Resten 1-113 der SEQ-ID-NR. 2 entsprechende Aminosäure-Sequenz kodiert.
4. Nucleinsäure nach Anspruch 1, welche die den Resten 1-125 der SEQ-ID-NR. 2 entsprechende Aminosäure-Sequenz kodiert.
5. Nucleinsäure nach Anspruch 1, welche die in SEQ-ID-NR. 2 dargestellte Aminosäure-Sequenz kodiert.
6. Isolierte Nucleinsäure, welche eine die durch SEQ-ID-NR. 2 beschriebene Transmembran-Domäne des HB15-Proteins kodierende Sequenz enthält.
7. Nucleinsäure nach Anspruch 6, welche die den Resten 126-147 der SEQ-ID-NR. 2 entsprechende Aminosäure-Sequenz kodiert.
8. Isolierte Nucleinsäure, welche eine die durch SEQ-ID-NR. 2 beschriebene cytoplasmatische Domäne des HB15-Proteins kodierende Sequenz enthält.
9. Nucleinsäure nach Anspruch 8, welche die den Resten 148-186 der SEQ-ID-NR. 2 entsprechende Aminosäure-Sequenz kodiert.
10. Isolierte Nucleinsäure, die eine ein Säugerhomologes des HB15-Proteins kodierende Sequenz enthält, wobei die Nucleinsäure unter stringenten Bedingungen mit einer DNA-Sonde, welche die in SEQ-ID-NR. 1 angegebene kodierende Sequenz enthält, hybridisieren kann und das Homologe die Gewebeverteilung hat, die für das durch die SEQ-ID-NR. 2 beschriebene menschliche HB15-Protein festgestellt wird.
11. Isolierte Nucleinsäure, die den Teil der Nucleinsäure nach Anspruch 10 enthält, der die extrazelluläre Domäne des Homologen kodiert.
12. Isolierte Nucleinsäure, die den Teil der Nucleinsäure nach Anspruch 10 enthält, der die Transmembran-Domäne des Homologen kodiert.
13. Isolierte Nucleinsäure, die den Teil der Nucleinsäure nach Anspruch 10 enthält, der die cytoplasmatische Domäne des Homologen kodiert.
14. Rekombinierter Vektor, der die Nucleinsäure nach einem der Ansprüche 1 bis 13 enthält.
15. Kultivierte Zellen, die mit dem Vektor nach Anspruch 14 transfiziert sind.
16. Zellen nach Anspruch 15, wobei die Zellen in der nicht-transfizierten Form das durch die Nucleinsäure kodierte Protein nicht exprimieren.
17. Verfahren zur Expression rekombinierter Nucleinsäure, bei dem die Zellen nach Anspruch 16 unter Bedingungen kultiviert werden, welche die Expression der Nucleinsäure, mit der sie transfiziert wurden, gestatten.
18. Verfahren zur Herstellung des durch SEQ-ID-NR. 2 beschriebenen HB15-Proteins, bei dem die kultivierten Zellen nach Anspruch 15 unter Bedingungen inkubiert werden, welche die Bildung des HB15-Proteins gestatten, und das HB15-Protein aus den inkubierten Zellen gewonnen wird.
19. Polynukleotid mit einer Länge von mehr als etwa 20 Nukleotiden, das unter stringenten Bedingungen an eine Nucleinsäure mit einer Sequenz, die komplementär zu einer in SEQ-ID-NR. 1 gezeigten Nucleinsäure-Sequenz ist, die das durch SEQ-ID-NR. 2 beschriebene HB15-Protein oder einen Teil davon mit dem Gewebeverteilungs-muster von HB15 kodiert, hybridisiert.
20. Polynukleotid nach Anspruch 19, wobei die stringenten Bedingungen eine 20% Formamid, 150 mM NaCl, 15 mM Trinatriumcitrat, 50 mM Natriumphosphat, 10% Dextranulfat und 20 µg/ml denaturierte, gescherte Lachssperma-

DNA enthaltende Hybridisierungslösung umfassen.

21. Polynukleotid nach Anspruch 19, das größer als etwa 50 Nukleotide ist.
- 5 22. Polynukleotid nach Anspruch 21, das größer als etwa 100 Nukleotide ist.
23. Polynukleotid nach Anspruch 19, wobei der Teil eine extrazelluläre Domäne des durch SEQ-ID-NR. 2 beschriebenen HB15-Proteins umfaßt.
- 10 24. Polynukleotid nach Anspruch 23, wobei der Teil die Aminosäuren 1-113 von SEQ-ID-NR.:2 umfaßt.
- 25 25. Kultivierte Zellen, die mit dem Polynukleotid nach Anspruch 19, das betriebsbereit an eine Transkriptions-Kontrollsequenz gebunden ist, transkribiert sind.
- 15 26. Zellen nach Anspruch 25, wobei die Zellen in der nichttranskribierten Form das durch das Polynukleotid kodierte Protein nicht exprimieren.
- 20 27. Verfahren zur Herstellung des durch das Polynukleotid nach Anspruch 19 kodierten Polypeptids, bei dem die Zellen nach Anspruch 25 unter zur Bildung des Polypeptids wirksamen Bedingungen kultiviert werden und das Polynukleotid gewonnen wird.
28. Polypeptid, das durch die Nukleinsäure nach einem der Ansprüche 1 bis 13 und 19 bis 24 kodiert wird.
29. Polypeptid mit einer Aminosäure-Sequenz des Polypeptids nach Anspruch 28.
- 25 30. Verfahren zur Isolation einer ein Homologes des durch SEQ-ID-Nr.:2 beschriebenen menschlichen HB15-Proteins kodierenden Nukleinsäure, bei dem die Nukleinsäure nach Anspruch 1 mit einer Population von Nukleinsäuremolekülen, die vermutlich ein ein HB15-Homologes kodierendes Nukleinsäuremolekül enthält, unter Hybridisierungsbedingungen hybridisiert wird, die zur Identifizierung des das HB15-Homologe kodierenden Nukleinsäuremoleküls in der Population durch Eintreten von Hybridisierung ausreichen, und daß das ein HB15-Homologes kodierende Nukleinsäuremolekül aus kreuzhybridisierten Molekülen isoliert wird.
31. Verfahren nach Anspruch 30, wobei die Population von Nukleinsäuremolekülen aus Mausgewebe erhalten wird.
- 35 32. Nukleinsäure-Isolat, das unter stringenten Hybridisierungsbedingungen an eine nach Anspruch 1 definierte Nukleinsäure hybridisiert.
33. Polypeptid, das durch die Nukleinsäure nach Anspruch 32 kodiert wird und das Gewebeverteilungsmuster des durch SEQ-ID-NR.:2 beschriebenen HB15-Proteins hat.
- 40 34. Polypeptid mit der Sequenz des Polypeptids nach Anspruch 33.
35. Monoklonaler Antikörper, der an das Polypeptid nach Anspruch 29 oder 34 bindet.
- 45 36. Monoklonaler Antikörper, der an das durch SEQ-ID-NR.:2 beschriebene HB15-Protein bindet.
37. Monoklonaler Antikörper, der an das HB15-Epitop bindet, das von einem Antikörper erkannt wird, der von der unter der ATCC-Nr.: HB10987 hinterlegten Zelllinie gebildet wird.
- 50 38. Monoklonaler Antikörper, der an das HB15-Epitop bindet, das von einem Antikörper erkannt wird, der von der unter der ATCC-Nr.: HB10988 hinterlegten Zelllinie gebildet wird.
39. Monoklonaler Antikörper, der von der unter der ATCC-Nr.: HB10987 hinterlegten Zelllinie gebildet wird.
- 55 40. Monoklonaler Antikörper, der von der unter der ATCC-Nr.: HB10988 hinterlegten Zelllinie gebildet wird.
41. Hybridom-Zelllinie, unter der ATCC-Nr.: HB10987 hinterlegt.

42. Hybridom-Zelllinie, unter der ATCC-Nr.: HB10988 hinterlegt.

43. Verfahren zur Isolierung von HB15 exprimierenden Zellen, bei dem der Antikörper nach einem der Ansprüche 35 bis 40 mit einer Population von Zellen umgesetzt und Zellen isoliert werden, an die der Antikörper bindet.

5 44. Verfahren zur Quantifizierung der von einer Population von Zellen exprimierten Menge an Oberflächen-HB15, bei dem ein Antikörper, der an das durch die SEQ-ID-NR. 2 beschriebene HB15-Protein bindet, mit einer Population von Zellen, von denen zumindest einige vermutlich Oberflächen-HB15 tragen, unter Bedingungen umgesetzt wird, welche die Bindung des Antikörpers an Oberflächen-HB15 gestalten, Zellen, an die der Antikörper bindet, nachgewiesen werden und die Menge an gebundenem Antikörper quantifiziert wird.

10 45. Monoklonaler Antikörper nach Anspruch 35, wobei der monoklonale Antikörper ein menschlicher monoklonaler Antikörper ist.

15 46. Monoklonaler Antikörper nach Anspruch 35, wobei der monoklonale Antikörper ein chimärischer monoklonaler Antikörper ist, der in der variablen Region von einem monoklonalen Maus-Antikörper stammende Segmente und von einem menschlichen Antikörper stammende andere Regionen aufweist.

20 **Revendications**

1. Acide nucléique isolé comprenant une séquence codant pour le domaine extracellulaire de la protéine HB15 décrite par SEQ ID N° : 2.
2. Acide nucléique selon la revendication 1, comprenant une séquence codant pour la séquence de HB15 entière présentée dans SEQ ID N° : 2.
3. Acide nucléique selon la revendication 1, codant pour la séquence d'aminoacides correspondant aux résidus 1-113 de SEQ ID N° : 2.
- 30 4. Acide nucléique selon la revendication 1, codant pour la séquence d'aminoacides correspondant aux résidus 1-125 de SEQ ID N° : 2.
- 5 5. Acide nucléique selon la revendication 1, codant pour la séquence d'aminoacides présentée dans SEQ ID N° : 2.
6. Acide nucléique isolé comprenant une séquence codant pour le domaine transmembranaire de la protéine HB15 décrite par SEQ ID N° : 2.
- 40 7. Acide nucléique selon la revendication 6, codant pour la séquence d'aminoacides correspondant aux résidus 126-147 de SEQ ID N° : 2.
8. Acide nucléique isolé comprenant une séquence codant pour le domaine cytoplasmique de la protéine HB15 décrite par SEQ ID N° : 2.
- 45 9. Acide nucléique selon la revendication 8, codant pour la séquence d'aminoacides correspondant aux résidus 148-186 de SEQ ID N° : 2.
10. Acide nucléique isolé comprenant une séquence codant pour un homologue de mammifère de la protéine HB15, ledit acide nucléique étant capable de s'hybrider dans des conditions stringentes à une sonde d'ADN comprenant la séquence codante présentée dans SEQ ID N° : 1, et ledit homologue ayant la distribution tissulaire observée pour la protéine HB15 humaine décrite par SEQ ID N° : 2.
- 50 11. Acide nucléique isolé comprenant la portion d'acide nucléique de la revendication 10 qui code pour le domaine extracellulaire dudit homologue.
12. Acide nucléique isolé comprenant la portion d'acide nucléique de la revendication 10 qui code pour le domaine transmembranaire dudit homologue.

13. Acide nucléique isolé comprenant la portion d'acide nucléique de la revendication 10 qui code pour le domaine cytoplasmique dudit homologue.

5 14. Vecteur recombiné comprenant l'acide nucléique de l'une quelconque des revendications 1 à 13.

15. Cellule cultivée transfectée avec le vecteur de la revendication 14.

16. Cellule selon la revendication 15, ladite cellule sous forme non transfectée n'exprimant pas la protéine codée par l'édit acide nucléique.

17. Procédé d'expression d'un acide nucléique recombiné, qui comprend la culture de la cellule selon la revendication 16 dans des conditions permettant l'expression de l'acide nucléique avec lequel elle a été transfectée.

18. Procédé de production de la protéine HB15 décrite par SEQ ID N° : 2, comprenant l'incubation de la cellule cultivée de la revendication 15 dans des conditions permettant la production de la protéine HB15, et la récupération de la protéine HB15 à partir de la cellule incubée.

19. Polynucléotide d'une longueur supérieure à environ 20 nucléotides, capable de s'hybrider dans des conditions stricte à un acide nucléique ayant une séquence complémentaire d'une séquence d'acide nucléique présente dans SEQ ID N° : 1 codant pour la protéine HB15 décrite par SEQ ID N° : 2 ou une de ses portions ayant le schéma de distribution tissulaire de HB15.

20. Polynucléotide selon la revendication 19, où les conditions strictes comprennent une solution d'hybridation contenant 20 % de formamide, du NaCl 150 mM, du citrate trisodique 15 mM, du phosphate de sodium 50 mM, 10 % de sulfate de dextrans et 20 µg/ml d'ADN de sperme de saumon découpé et dénaturé.

21. Polynucléotide selon la revendication 19, qui a une longueur supérieure à environ 50 nucléotides.

22. Polynucléotide selon la revendication 21, qui a une longueur supérieure à environ 100 nucléotides.

23. Polynucléotide selon la revendication 19, dans lequel ladite portion comprend un domaine extracellulaire de la protéine HB15 décrite par SEQ ID N° : 2.

24. Polynucléotide selon la revendication 23, dans lequel ladite portion comprend les aminoacides 1 à 113 de SEQ ID N° : 2.

25. Cellule cultivée, transfectée avec le polynucléotide de la revendication 19 lié de manière opérationnelle à une séquence de contrôle de la transcription.

40 26. Cellule selon la revendication 25, ladite cellule sous forme non transfectée n'exprimant pas la protéine codée par l'édit polynucléotide.

45 27. Procédé de production du polypeptide codé par le polynucléotide de la revendication 19, comprenant la culture des cellules de la revendication 25 dans des conditions efficaces pour la production dudit polypeptide, et la récupération dudit polypeptide.

28. Polypeptide codé par l'acide nucléique de l'une quelconque des revendications 1 à 13 et 19 à 24.

50 29. Polypeptide ayant une séquence d'aminoacides du polypeptide de la revendication 28.

30. Procédé d'isolement d'un acide nucléique codant pour un homologue de la protéine HB15 humaine décrite par SEQ ID N° : 2, comprenant l'hybridation de l'acide nucléique de la revendication 1 avec une population de molécules d'acides nucléiques suspectée de contenir une molécule d'acide nucléique codant pour un homologue de HB15 dans des conditions d'hybridation suffisantes pour l'identification de ladite molécule d'acide nucléique codant pour l'homologue de HB15 dans ladite population par la production éventuelle d'une hybridation ; et l'isolement de ladite molécule d'acide nucléique codant pour l'édit homologue de HB15 à partir des molécules formées par hybridation croisée.

31. Procédé selon la revendication 30, dans lequel ladite population de molécules d'acide nucléique est obtenue à partir de tissu de souris.

32. Isolat d'acide nucléique capable de s'hybrider dans des conditions d'hybridation stringentes à un acide nucléique tel que défini par la revendication 1.

33. Polypeptide codé par l'acide nucléique de la revendication 32 et ayant le schéma de distribution tissulaire de la protéine HB15 décrite par SEQ ID N° : 2.

34. Polypeptide ayant la séquence du polypeptide de la revendication 33.

35. Anticorps monoclonal qui se lie au polypeptide de la revendication 29 ou 34.

36. Anticorps monoclonal qui se lie à la protéine HB15 décrite par SEQ ID N° : 2.

37. Anticorps monoclonal qui se lie à l'épitope de HB15 reconnu par un anticorps produit par la lignée cellulaire déposée sous le n° ATCC HB10987.

38. Anticorps monoclonal qui se lie à l'épitope de HB15 reconnu par un anticorps produit par la lignée cellulaire déposée sous le n° ATCC HB10988.

39. Anticorps monoclonal produit par la lignée cellulaire déposée sous le n° ATCC HB10987.

40. Anticorps monoclonal produit par la lignée cellulaire déposée sous le n° ATCC HB10988.

41. Lignée cellulaire d'hybridome déposée sous le n° ATCC HB10987.

42. Lignée cellulaire d'hybridome déposée sous le n° ATCC HB10988.

43. Procédé d'isolement de cellules exprimant HB15, comprenant la réaction de l'anticorps de l'une quelconque des revendications 35 à 40 avec une population de cellules, et l'isolement des cellules auxquelles se lie ledit anticorps.

44. Procédé de détermination de la quantité de HB15 de surface exprimé sur une population de cellules, comprenant la réaction d'un anticorps qui se lie à la protéine HB15 décrite par SEQ ID N° : 2 avec une population de cellules dont au moins certaines sont suspectées de porter la HB15 de surface, dans des conditions qui permettent audit anticorps de se lier à la HB15 de surface; la détection des cellules auxquelles se lie ledit anticorps; et la détermination de la quantité d'anticorps lié.

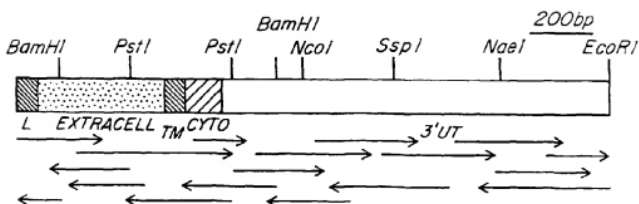
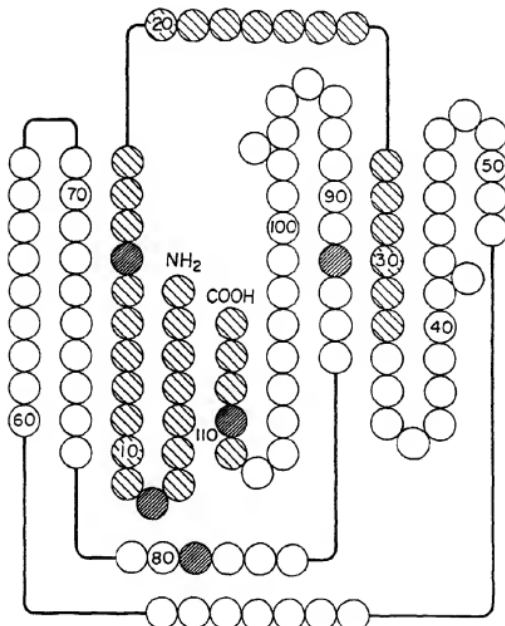
45. Anticorps monoclonal selon la revendication 35, l'anticorps monoclonal étant un anticorps monoclonal humain.

46. Anticorps monoclonal selon la revendication 35, l'anticorps monoclonal étant un anticorps monoclonal chimère ayant des segments de régions variables dérivés d'un anticorps monoclonal de souris et d'autres régions dérivées d'un anticorps humain.

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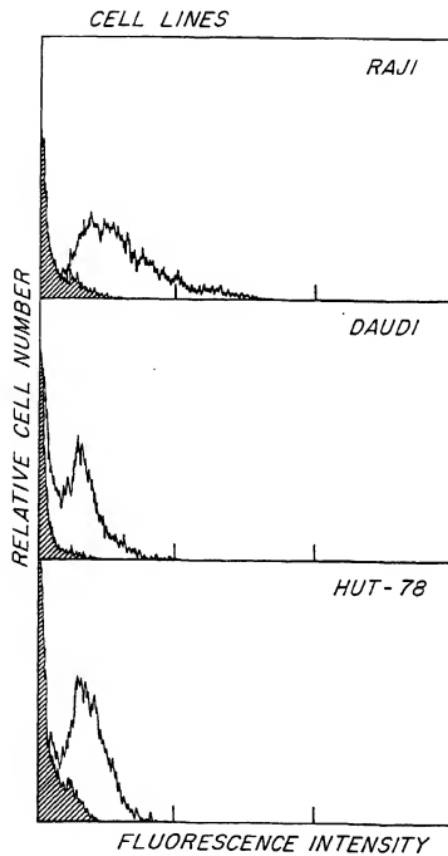
**FIG. 1****FIG. 3**

gaattccGCC M S R G L Q L L S C A Y S L A  
ATG TCC CGC GGC CTC CAG CTT CTG CTC CTC 61  
 P A T 1 P E V K V A C O S E D V D L P C T A  
CCG CGG ACG CCG GAG GTG AAG GTG GCT TG  
 + 20 + TCC TGA GAT GTG GAC TTG CCC TGC ACC GCC 121  
 P W D P Q V P Y T V S W V K L L E G G E  
CCC TGG GAT CCG CAG GGT CCC TAC ACG GTC TCC  
 40 50 A GTC AAC TTA TTG GTC AAC GGT GAA 181  
 E R M E T P Q E D H L R G Q H Y H Q K G  
GAG AGG ATG GAG ACA CCC CAG GAA GAC CAC CTC  
 60 70 + AGG CAC TCA CAC TAT CAT CAG AAG GGG 241  
Q N G S F D A P N E R P Y S L K I R N T  
CAA AAT GGT TCT TTC GAC GCC CCC AAT GAA AGG  
 + 80 + CCC TAT TCC CTC AAG ATC CGA AAC ACT 301  
T S C O N S G T Y R C O T L Q D P D G Q R N  
ACC AGC TGC AAC TCG GGG ACA TAC AGG TGC ACT  
 100 110 CTC GAC CGG GAT GGG CAG AGA AAC 361  
L S G K V I L R V T G C O P A Q R K E E T  
CTA AGT GGC AAG GTG ATC TTG AGA GTG ACA GGA  
 120 130 TGC CCT GCA CAG CGT AAA GAA GAG ACT 421  
 F K K Y R A E I V L L A L V I F Y L T  
TTT AAG AAA TAC AGA GCG GAG ATT GTC CTC GCT  
 140 150 GCT ATT TTC ACT TGT(RAG)TTT GCA CGG CTA CAG AGT ATC TTC CCA GAT TTT TCT AAA 481  
 L I I F T C O K F A R L Q S I F P D F S K  
CTC ATC ATT TTC ACT TGT(RAG)TTT GCA CGG CTA  
 160 170 GCT ACC TCC CCA GTC ATT AAT AAC CAT TTA GGG CTA GTG 541  
 A G M E R A F L P V T S P N K H L G L V  
GCT GGC ATG GAA CGA GCT TTT CTC 601

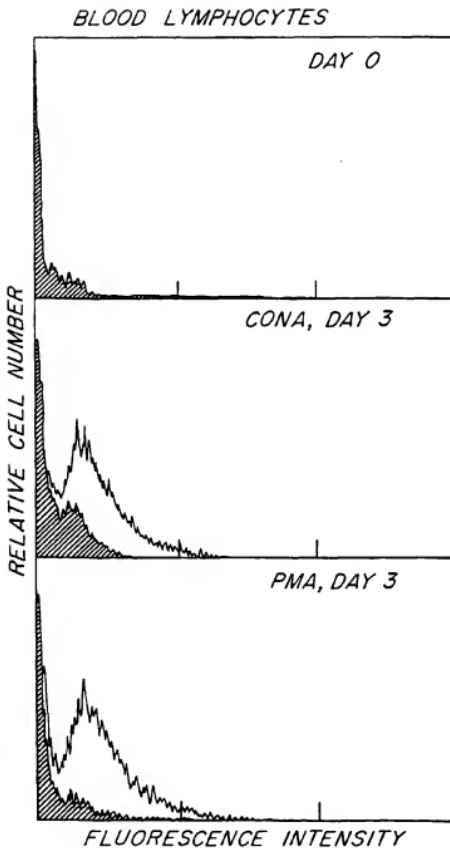
*FIG. 2A*

180	T	P	H	K	T	E	L	*		
	ACT	CCT	CAC	AAG	ACA	GAA	CTG	GTA	TGA	GCAGGGATTIC
	AGGGGTGTC	CTGCTGTTA	CACMGGAGGA	GAGAAGANTG	AGCCATGGCT	GAAGATGCCA	TCCCTGTGAG			GTCTGAGGCTC
	TCTTCACCT	CACTGAAAC	ATCGGGAAAG	GGATCCACC	CCATTTCTG	TGGCAGGGCC	TGAAAAACCA			668
	TCACATGACC	ACATAGCATG	AGGCCACATG	TGCTTCTCCA	TGGCACCTT	TGCGCACCTT	TGAAAGGCTA			738
	TCTGGTCAAC	CTCTCTGACA	TITTTTCACT	CATAAAAGG	CTATGGGAG	ATGCACTGG	AAAGGGCTA			808
	TGGGAAATATC	GAATGCCCCC	AGCTGGCCCG	TGCAAGACTC	CTGAGGACAG	CTGCACTCTG				878
	GGGACATTC	TITTGATTTT	CTGTGTTTTC	CGTGATCCAG	CCAGAAGTTT	TACGCTGGG				948
	AGATCAGCT	GTGAGACAGT	GGAAATAATT	TAGCAAAATA	TITTCCTGTT	TGAAGGTCCT	AGRAATTGAC			1018
	AGGAGTAACTC	TGTGTACAAA	GAATAACAA	GTGCGATGAC	TATTCCCCAG	CAGEGCTCTT	GCATATTACTA			1088
	AAGACATCCA	TAANGAAGCA	ATTAAGAAGA	GTGCCACATT	TATTTCATTA	TCATCTGGGA				1158
	AAGGTTGGT	TTTTCTGCT	TTTGAATCT	TCATTTATTA	TCTATAGTA	CTTGTCAAG				1228
	TGGACATAGA	GAEGGGAGAG	AACTCAGAGA	GGGTGACAG	TTGCAACTG	ACAGGCAGCC				1298
	TGCCGGGCGT	ACGGTGCACT	CTGGGGTCT	GTCCACTGT	ATTAATGGC	CGGTGCGRAA				1368
	GTTCCTTCG	GTGCTGCTT	TCCTCCATGT	GTCCACTGT	CCCACTATCT	GGGTGCAATG				1438
	CGTGAAATAAT	CTTGTAAAC	ATATAAACTA	GTGTGTGCA	GGGGCTAATGA	TCTTGAGATA				1508
	ATGATGATGG	TGTTTGGG	ATGAGAGGGT	GCTATCCATT	GCTGTGCA	AGATCATATA				1578
	AGGTTACCAA	GAAGCCCTTC	CTGTAGCCCT	TCTCATGTTT	TCCATTGTTT	GAACAAAGA				1648
										1718
										1761

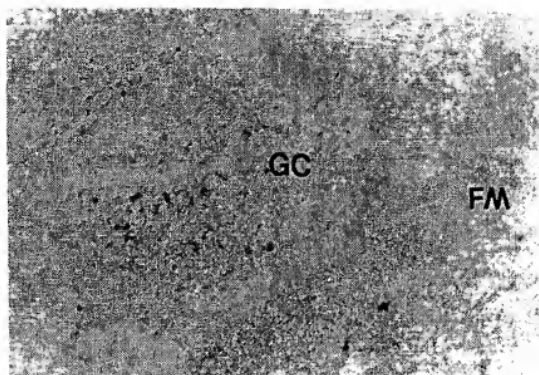
## FIG. 2B



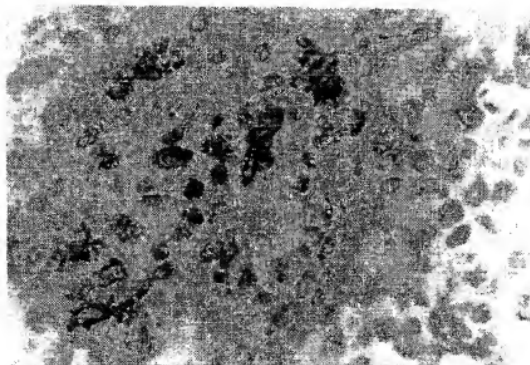
**FIG. 4A**



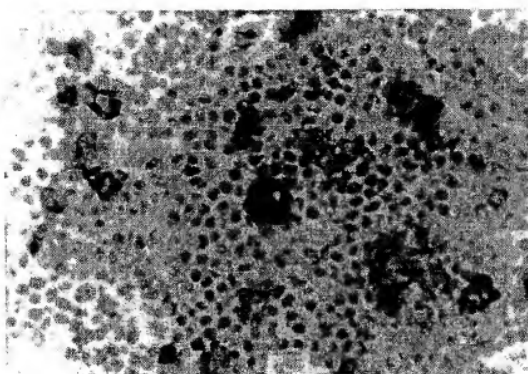
**FIG. 4B**



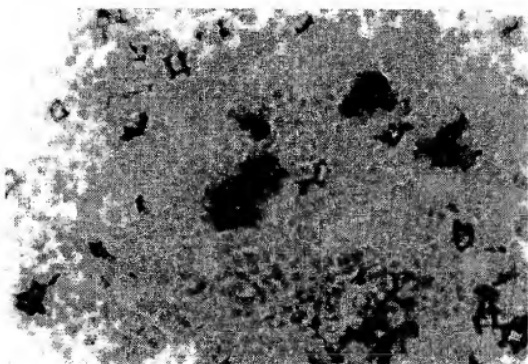
*FIG. 5A*



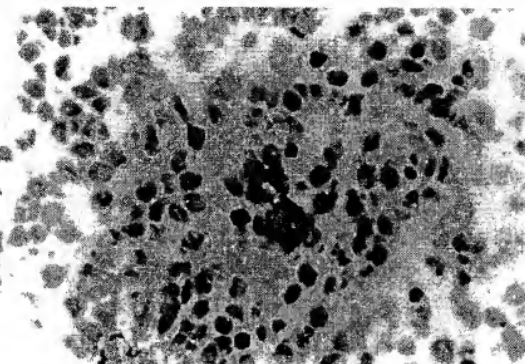
*FIG. 5B*



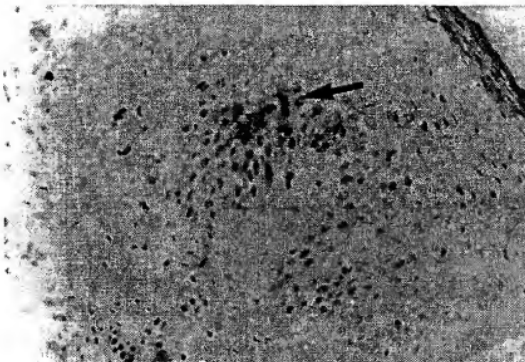
*FIG. 5C*



*FIG. 5D*



*FIG. 5E*



*FIG. 5F*